PARASITISM OF A FACTITIOUS HOST, GALLERIA MELLONELLA (LEPIDOPTERA: PYRALIDAE) BY AN ENDOPARASITOID: OVIPOSITION AND EMERGENCE OF MICROPLITIS CROCEIPES (HYMENOPTERA: BRACONIDAE)

PREM GUPTA, ALEXIS SLOAN, CHARLES R. DILLARD, AND STEPHEN M. FERKOVICH Insect Attractants, Behavior, and Basic Biology Research Laboratory, Agricultural Research Service, U.S. Department of Agriculture, Gainesville, FL 32604

ABSTRACT

The effect of various diet supplements on the development of *Microplitis croceipes* in an atypical host, *Galleria mellonella* (Linnaeus), were evaluated, as were ovipositional responses to various factors. Female parasitoids were exposed to fifth instar *G. mellonella* in Petri dishes containing the following treatments either separately or in combination: a) *Helicoverpa zea* (*Boddie*) frass, b) 1% and 10% solutions of a host-seeking stimulant (13-methylhentriacontane), c) *H. zea* hemolymph, and d) *H. zea* hemolymph concentrated by freeze-drying. There were no significant differences be-

tween hemolymph and frass + hemolymph treatments. The host-seeking stimulant alone also stimulated oviposition. The most effective combination was host-seeking stimulant and concentrated hemolymph which induced oviposition rates comparable to that in the typical host, H. zea. Various diet supplements did not improve the development and emergence of M. croceipes. We conclude that oviposition by M. croceipes in the atypical host, G. mellonella, was significantly improved by the application of host-seeking stimulant and concentrated hemolymph, but the rate of adult parasitoid emergence was not increased by the addition of nutrient supplements to the host diet.

Key Words: Rearing, in vivo, host-seeking stimulant, diet supplement.

RESUMEN

Fueron evaluados el efecto de varios suplementos en la dieta sobre el desarrollo de Microplitis croceipes en un hospedante atípico, Galleria mellonella (Linnaeus), y la respuesta ovoposicional a varios factores. Las hembras del parasitoide fueron expuestas al quinto estadio de G. mellonella en placas de Petri conteniendo los siguientes tratamientos, separados o combinados: a) excretas de Helicoverpa zea (Boddie), b) soluciones de estimulante de búsqueda del hospedante (13-methylhentriacontane), c) hemolinfa de H. zea, y d) hemolinfa de H. zea concentrada mediante secado por congelación. No hubo diferencias significativas entre los tratamientos de hemolinfa y hemolinfa-excretas. El estimulante solo también aumentó la ovoposición. La combinación más efectiva fue estimulante con hemolinfa concentrada que estimuló la ovoposición hasta valores comparables a la tasa de ovoposición en el hospedante típico, H. zea. Varios suplementos a la dieta del hospedante no mejoraron el desarrollo y la emrgencia de M. croceipes. Concluimos que la ovoposición por M. croceipes en el hospedante atípico, G. mellonella, fue significativamente mejorada mediante la aplicación de estimulante de búsqueda y hemolinfa concentrada, pero la tasa de emergencia del parasitoide adulto no fue incrementada mediante la adición de nutrientes suplementarios a la dieta del hospedante.

Microplitis croceipes (Cresson) is a solitary endoparasitoid that attacks the corn earworm, Helicoverpa zea (Boddie) and tobacco budworm, Heliothis virescens (Fabricius). The mass rearing of M. croceipes is important because of its potential use in biological control of Heliothis/Helicoverpa species (Knipling & Stadelbacher 1983). However, mass rearing of M. croceipes is expensive because host larvae are cannibalistic and must be reared individually (Powell & Hartley 1987; Greany et al. 1989; King & Coleman 1989). A large number of insect parasitoids have been successfully cultured in vitro (Grenier et al. 1994), but none of the hymenopteran larval endoparasitoids have been reared from egg to adult in vitro (Thompson 1986; Rotundo et al. 1988; Greany, 1991; Ferkovich & Oberlander 1991; Pennachio et al. 1992; Ferkovich et al. 1994). We studied six species of Lepidoptera as potential atypical hosts for in vivo rearing of M. croceipes, but successful development of M. croceipes occurred only in G. mellonella (21% adult emergence) and to a lesser extent in the fall armyworm Spodoptera frugiperda (13% adult emergence) (Blumberg & Ferkovich 1994; Ferkovich & Blumberg 1994). The advantage of using Galleria mellonella as a factitious host for rearing is that it is often less expensive to rear than other species. For example, the cost of rearing the tachinid parasitoid Lixophaga diatraea was reduced 81% when reared on G. mellonella as compared with costs of production on the natural host, Diatraea saccharalis (King et al. 1979). Estimated rearing costs for diet, labor and containers in our laboratory for *G. mellonella* are 3X less costly than for *H. zea*, and automated large-scale rearing of *G. mellonella* is expected to be even less expensive. However, for practical mass rearing, the rates of atypical-host parasitism and parasitoid adult eclosion from the atypical host must be improved (Gupta et al. 1996). Here, we focus on (1) improvement of the rate of oviposition, and (2) the effect of some nutritional supplements (not routinely present in the diet of *G. mellonella*) as they relate to development and emergence of *M. croceipes*.

MATERIALS AND METHODS

Parasite-Host Colony Maintenance

H. zea was reared according to Lewis & Burton (1979) at the USDA/ARS, Insect Biology and Population Management Research Laboratory, Tifton, Georgia. We received eggs by mail and allowed them to hatch on Heliothis Premix diet® (Stonefly Industries, Inc., Bryan, Texas). M. croceipes was reared as described by Ferkovich & Dillard (1986) with the following modifications. Two 3-day-old female parasitoids were added to each cup of 40-50 H. zea late second and early third instars for 24 h. Larvae were then removed, individually placed in 1 ounce plastic cups with 4-5 ml of semisolid diet and kept at 25°C and 60-70% relative humidity for 14 days. Parasitoid cocoons were removed and placed in an emergence cage at 26°C, 55% relative humidity and a photoperiod of 15:9 [L:D] for three weeks. Parasitoid adults were held at a sex ratio of 1:1 in Plexiglas cages streaked with honey. At 2 days, males were discarded and females kept for colony maintenance and research. G. mellonella was reared according to Bean & Silhacek (1989).

Experimental

Fifth instar G. mellonella larvae (28 mg avg. wt.) not previously exposed to parasitoids were placed in Petri dishes (9 cm diam) with 4-d-old females in the ratio of 2:1 (host:parasitoid) for 1 h. Fifth instar G. mellonella were used because they were similar in size to third instar H. zea which parasitoid females readily attacked. Fifty to 100 larvae of G. mellonella (5 to 10 replicates of 10 larvae each) were stung for the various treatments. Oviposition by M. croceipes was determined by counting the first instar parasitoids dissected from host larvae three days after parasitization. Frass and host-seeking stimulant solutions (10 μ l) were smeared on the bottom of the Petri dish, leaving a coat of the material on the surface. Hemolymph was collected from early fourth instars of H. zea by clipping a proleg. Each G. mellonella larva was then rolled in a drop of hemolymph and immediately exposed to female wasps. To prevent melanization while concentrating the hemolymph, 10 μ l of 5% solution of phenylthiourea in methanol was added per 500 μ l of hemolymph held on ice. Hemolymph was concentrated two-fold using a lyophilizer (Virtis Research Equipment, Gardiner, New York).

Because untreated G. mellonella host larvae do not elicit an ovipositional response by M. croceipes females (Ferkovich & Blumberg 1994), they were given the following treatments just before exposing them to M. croceipes females: 1) H. zea frass at the bottom of the Petri dish; 2) H. zea hemolymph; 3) frass + H. zea hemolymph; 4) two-fold concentrated freeze-dried H. zea hemolymph; 5) 1% host-seeking stimulant (13-methylhentriacontane); 6) 10% host-seeking stimulant; 7) 1% host-seeking stimulant plus two-fold concentrated freeze-dried H. zea hemolymph; 8) 10% host-seeking stimulant plus two-fold concentrated freeze-dried H. zea hemolymph. The host-seeking stimulant, purified from the feces and larvae of H. zea, triggers the short-range host

seeking response of M. croceipes females. Host-seeking stimulant was provided by Dr. R. E. Doolittle, USDA/ARS, Gainesville, Florida. This compound was dissolved in hexane and used at a concentration of 1% and 10% ($10\,\mu$ l) at the bottom of the Petri plate.

The following agents were added to the standard diet of fifth instar G. mellonella immediately after parasitization by M. croceipes: fetal bovine serum, chicken serum, powdered Grace's medium and TC-100 powdered medium (purchased from Gibco BRL, Grand Island, New York). The following nutrient supplements were also added: 1) torula dried yeast, 5% (Rhinelander, Wisconsin); 2) powdered whole liver, 5% (Schiff, Salt Lake City, Utah); 3) H. zea hemolymph, 2.5%; 4) fetal bovine serum, 5%; 5) chicken serum, 5%; 6) egg yolk, 5% (Sonstegard Food Co, Sioux Falls, South Dakota); 7) powdered Grace's medium at 10, 20, and 50 mg/gm of fresh diet; 8) powdered 52-B medium at 20 and 50 mg/gm of fresh diet (JRH Biosciences, Lenexa, Kansas); 9) powdered TC-100 at 10 mg/gm; 10) 50:50 mixture of Burton's modified pinto bean diet used in rearing of H. zea. (Burton & Perkins 1972, Milton's Institutional Foods Inc., Oakwood, Georgia) and G. mellonella diet (Bean & Silhack 1989). Fetal bovine serum, chicken serum and H. zea hemolymph were lyophilized before they were added to the dry components of the G. mellonella diet. All supplements were mixed thoroughly with the dry components of the G. mellonella diet prior to the addition of water-honeyglycerol. Diet supplement concentrations were based on the final fresh weight of the diet.

Statistical Analysis

Data were analyzed by one-way analysis of variance (ANOVA), and means were separated by the Tukey-Kramer multiple comparisons test using Instat II $^{\circ}$ (Graphpad Software, San Diego, California). All significance reported is at the P=0.05 level.

RESULTS

Concentrated Hemolymph and Host-Seeking Stimulant

The combination of frass plus hemolymph did not significantly improve percent parasitism compared to frass or hemolymph tested alone (Fig. 1). One percent and 10% host-seeking stimulant applications improved parasitism over frass, hemolymph, or hemolymph plus frass, although the higher dose (10%) was not statistically significant due to the large variation among replicates. Concentrated hemolymph significantly increased parasitization compared to that induced by frass, hemolymph, or frass plus hemolymph and was comparable to the two host-seeking stimulant treatments. The 10% concentration of host-seeking stimulant was not significantly different from the 1% concentration. However, when 1 and 10% concentrations of host-seeking stimulant were applied in combination with 2× concentrated hemolymph, the rate of parasitism was greater than all the other treatments and, more importantly, these values were comparable to ovipositional rate in the typical host *H. zea.*

Nutritional Supplements

Of all the diet supplements tested, only Grace's tissue culture medium (20 mg/gm) significantly enhanced percent cocoon production relative to the control diet (22.7% vs 18.49%). None of the other diet supplements tested significantly improved the development and/or emergence of M. croceipes adults (Table 1).

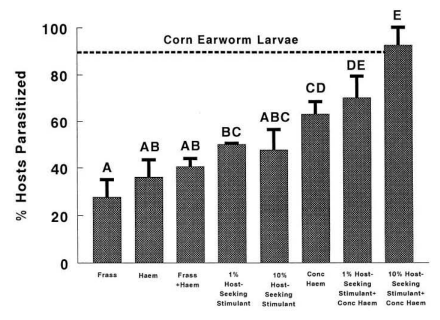


Figure 1. Effects of eight treatments on the percent parasitism of the atypical host Galleria mellonella, by Microplitis croceipes. All values are means \pm SE. Mean followed by the same letter are not significantly different (P> 0.05).

DISCUSSION

Parasitoids often do not oviposit in atypical hosts because chemical and physical stimuli associated with their usual host are absent. However, oviposition may be increased by (1) the application of contact chemicals extracted from the host frass and (2) secretions of the host mandibular and silk glands which can be perceived by the parasitoids (Vinson 1975; Waage et al. 1985). Recently, we demonstrated the acceptance of six atypical lepidopterans as candidate hosts for rearing M. croceipes (Ferkovich & Blumberg 1994). These atypical hosts were acceptable for oviposition after treatment with H. zea hemolymph or frass or the combination of frass and hemolymph. Host-seeking stimulant, which accentuates host seeking in M. croceipes (Jones et al. 1971), is more effective in combination with hemolymph and improves oviposition in G. mellonella. Previous observations on oviposition by M. croceipes females indicated that they responded to a two-component ovipositional kairomone in the hemolymph of the host (Tilden & Ferkovich 1988; Eller et al. 1990; Heath et al. 1990). Our results clearly show that the application of host-seeking stimulant, in combination with two-fold concentrated H. zea hemolymph, stimulated oviposition by M. croceipes in the atypical host, G. mellonella, at a rate comparable to oviposition in its natural host, H. zea.

Achieving a rate of oviposition in a factitious host that is comparable to that in its natural host is an important factor in effective rearing of the parasitoid. In future studies, application of purified kairomone from hemolymph of *H. zea* may be possible (Heath et al. 1990; Drost & Carde 1992).

Increased oviposition does not by itself denote successful parasitization. Thus, we studied the effect of the addition of a wide range of biologically active nutrient supple-

Table 1. Effects of different diet supplements on the development and emergence of M. CROCEIPES from G. MELLONELLA.

	Treatment	% Parasitism	% Cocoon	% Emergence	% Males	% Females
1.	Control	57.6 ^b	18.41 [™]	66.6ª	50.0ª	50.0 ^b
		(49/85)	(9/49)	(6/9)	(3/6)	(3/6)
2.	Grace's	59.1^{ab}	22.7^{a}	63.6ª	57.1^{a}	42.9^{b}
	20mg/gm	(97/164)	(22/97)	(14/22)	(8/14)	(6/14)
3.	Grace's	58.2 ^b	15.3ab	83.3°	40.0^{a}	60.0^{b}
	50mg/gm	(78/134)	(12/78)	(10/12)	(4/10)	(6/10)
4.	52-B	69.7ª	3.0°	100.0°	50.0^{a}	50.0^{a}
	20mg/gm	(62/89)	(2/62)	(2/2)	(1/2)	(1/2)
5 .	52-B	61.4^{ab}	9.2°	60.0ª	66.7^{a}	33.3^{a}
	50mg/gm	(54/88)	(5/54)	(3/5)	(1/3)	(2/3)
6.	50HPM3:50	41.7°	14.2be	40.0ª	0.0^{a}	100.0 ^b
	GPM Diet	(35/84)	(5/35)	(2/5)	(0/2)	(2/2)
1.	Control	62.8ª	18.4^{a}	44.4	25.0^{a}	75.0^{a}
		(49/88)	(9/49)	(4/9)	(1/4)	(3/4)
2.	Torula-	46.7ª	11.6ª	80.0ª	50.0^{a}	50.0^{a}
	Yeast	(43/92)	(5/11)	(4/5)	(2/4)	(2/4)
3.	H. Zea Hemo-	51.1°	10.9ª	80.0ª	50.0^{a}	50.0^{a}
	lymph~(2.5%)	(49/90)	(5/46)	(4/5)	(2/4)	(2/4)
1.	Control	77.5^{a}	18.8^{a}	76.9ª	40.0^{a}	60.0ª
		(69/89)	(13/69)	(10/13)	(4/10)	(6/10)
2.	TC-100	75.6ª	7.4^{b}	60.0^{a}	33.3^{a}	66.7^{a}
	10mg/gm	(68/90)	(5/68)	(3/5)	(1/3)	(2/3)
3.	Grace's	63.4^{a}	25.0^{a}	61.5^{*}	50.0^{a}	50.0^{a}
	10mg/gm	(52/82)	(13/52)	(8/13)	(4/8)	(4/8)
1.	Control	85.4^{a}	34.1^a	28.6^{a}	50.0ª	50.0^{a}
		(41/48)	(14/41)	(4/14)	(2/4)	(2/4)
2.	Whole	89.0ª	14.3^{a}	57.1°	25.0^{a}	75.0^{a}
	$\mathrm{Liver}(5\%)$	(49/55)	(7/49)	(4/7)	(1/4)	(3/4)
1.	Control	73.6ª	17.9^{a}	50.0ª	33.3°	66.7ª
		(67/91)	(12/67)	(6/12)	(2/6)	(4/6)

¹Diet supplement concentrations are on the basis of final fresh weight of the diet. Fifth instar G. mellonella were reared on these supplemented diets after parasitization. New controls were run with experiments that were run on different dates.

²Percentages within each column and each test group followed by the same letter are not significantly different (P > 0.05).

³HPM refers to 50% H. xea Premix diet and 50% G. mellonella diet.

Table 1. (Continued) Effects of different diet supplements 1 on the development and emergence of M. CROCEIPES from G. MELLONELLA. 2

	Treatment	% Parasitism	% Cocoon	% Emergence	% Males	% Females
2.	Fetal Bovine	51.4ª	8.1ª	33.3ª	100.0ª	0.0ª
	Serum (5%)	(37/72)	(3/37)	(1/3)	(1/1)	(0/1)
1.	Control	70.3ª	31.6ª	75.0ª	44.4ª	55.6°
		(38/54)	(12/38)	(9/12)	(4/9)	(5/9)
2.	Chicken	58.3ª	26.2^{a}	45.4^{a}	60.0^{a}	40.0^{a}
	Serum (5%)	(42/72)	(11/42)	(5/11)	(3/5)	(2/5)
1.	Control	58.4^{a}	28.4ª	53.8ª	42.8^{a}	57.2ª
		(45/77)	(13/45)	(7/13)	(3/7)	(4/7)
2.	Egg Yolk	56.9ª	29.7^{a}	45.4^{a}	40.0^{a}	60.0^{a}
	(5%)	(37/65)	(11/37)	(5/11)	(2/5)	(3/5)

'Diet supplement concentrations are on the basis of final fresh weight of the diet. Fifth instar G. mellonella were reared on these supplemented diets after parasitization. New controls were run with experiments that were run on different dates.

ments directly to the diet of G. mellonella to support the development of M. croceipes. However, none of the supplements tested significantly affected the development and emergence of M. croceipes.

The possibility of mass-rearing parasitoids for use in biological control programs is associated with the parasitoids' degree of interaction with the host's physiology, as suggested by Campadelli & Dindo (1988). The nutritional and ecological considerations in propagation of entomophagous and endoparasitic insects are evaluated in detail by Thompson (1990) and Slansky (1986). In hymenopteran endoparasitoids, physiological interactions are complex, and it seems from our earlier in vitro studies (Ferkovich et al. 1994) and from the present in vivo data, that it may not be feasible to manipulate the development and emergence of M. croceipes by supplemental nutrients. However, recent studies by Carpenter (personal communication; see Gross et al. 1995) show that addition of torula yeast to the G. mellonella diet increased the weight of mature G. mellonella larvae and also of males and females of the tachnid parasitoid Archytas marmoratus reared from these G. mellonella.

In conclusion, these data indicate that although oviposition by $M.\ croceipes$ in the atypical host $G.\ mellonella$ can be significantly improved by the application of host-seeking stimulant and two-fold concentrated hemolymph. The low emergence of adult parasitoids still remains a major problem for rearing the parasitoid on this host.

ACKNOWLEDGMENTS

The authors thank Dr. R. Ngugen, Dr. J. Sivinski, Dr. P. Greany, Dr. H. Oberlander, and P. Egger for their helpful comments.

Percentages within each column and each test group followed by the same letter are not significantly different (P > 0.05).

HPM refers to 50% H. zea Premix diet and 50% G. mellonella diet.

REFERENCES CITED

- BEAN, D. W., AND D. L. SILHACK. 1989. Changes in the titer of the female-predominant storage protein (81K) during larval and pupal development of the waxmoth, *Galleria mellonella*. Arch. Insect Biochem. & Physiol. 10: 333-348.
- Blumberg, D., and S. M. Ferkovich. 1994. Development and encapsulation of the endoparasitoid, *Microplitis croceipes* (Hymenoptera: Braconidae), in six candidate host species. Entomophaga 39: 293-302.
- BURTON, R. L., AND W. D. PERKINS. 1972. WSB, a new laboratory diet for the corn earworm and the fall armyworm. J. Econ. Entomol. 65: 385-386.
- CAMPADELLI, G., AND M. L. DINDO. 1988. Recent progress in the study of artificial diets for rearing insect parasitoids. Bollettino-dell'Instituto-di-Entomologia-della-Universita-delgi-Studi-di-Bologna. 42: 101-118.
- DROST, Y. C., AND C. R. CARDE. 1992. Host switching in *Brachymeria intermedia* (Hymenoptera: Chalcididae), a pupal endoparasitoid of *Lymantria dispar* (Lepidoptera: Lymantriidae). Environ. Entomol. 21: 760-766.
- ELLER, F. J., R. R. HEATH, AND S. M. FERKOVICH. 1990. Factors affecting oviposition by the parasitoid *Microplitis croceipes* (Hymenoptera: Braconidae). J. Econ. Entomol. 83: 398-404.
- FERKOVICH, S. M., AND C. R. DILLARD. 1986. A study of radiolabelled host proteins and protein synthesis during development of eggs of the endoparasitoid, *Microplitis croceipes* (Cresson) (Braconidae). Insect Biochem. 16: 337-345.
- FERKOVICH, S. M., AND H. OBERLANDER. 1991. Stimulation of endoparasitoid egg development by a fat body cell line: activity and characterization of factors that induce germ band formation and hatching, pp. 181-187 in Proceedings of the VIII International Conference on Invertebrate and Fish Tissue Culture. Tissue Culture Assoc., Columbia, MD.
- Ferkovich, S. M., H. Oberlander, C. R. Dillard, and C. E. Leach. 1994. Embryonic development of an endoparasitoid, *Microplitis croceipes* (Hymenoptera:Braconidae) in cell line-conditioned media. In Vitro Cell. Dev. Biol. 30: 279-282.
- FERKOVICH, S. M., AND D. BLUMBERG. 1994. Acceptance of six atypical host species for oviposition by *M. croceipes* (Hymenoptera: Braconidae). Israel J. Entomol. 28: 123-131.
- GREANY, P. D. 1991. Innovations in culturing cells, tissues, and endoparasites of invertebrates. In Vitro Cell Dev. Biol. 27A: 469.
- GREANY, P. D., S. M. FERKOVICH, AND W. R. CLARK. 1989. Progress toward development of an artificial diet and an in vitro rearing system for Microplitis croccipes. Southwest Entomol. 12: 89-94.
- GRENIER, S., P. D. GREANY, AND A. C. COHEN. 1994. Potential for mass release of insect parasitoids and predators through development of artificial culture techniques, pp. 181-205 in D. Rosen, F. D. Bennett and J. L. Capinera [eds.]. Pest management in the subtropics: Biological control-a Florida perspective. Intercept Publishers, Andover, Hampshire, England.
- GROSS, H. R., C. E. ROGERS, AND J. E. CARPENTER 1995. Development of *Archytas marmoratus* (Diptera: Tachinidae) reared in *Galleria mellonella* larvae (Lepidoptera: Pyralidae) feeding on selected diets. (unpublished data).
- GUPTA, P., C. R. DILLARD, AND S. M. FERKOVICH. 1996. Potential of an unnatural host, G.mellonella for rearing the corn earworm endoparasitoid Microplitis croceipes (Hymenoptera: Braconidae). Ann. Entomol. Soc. America 89: 103-108.
- HEATH, R. R., S. M. FERKOVICH, P. D. GREANY, F. J. ELLER, B. D. DUEBEN, AND R. L. TILDEN. 1990. Progress in the isolation and characterization of a host hemolymph ovipositional kairomone for the endoparasitoid *Microplitis croceipes*. Arch. Insect Biochem. Physiol. 13: 255-265.
- JONES R. L., W. J. LEWIS, M. BEROZA, AND B. A. BIERL. 1971. Host-seeking stimulant for parasite of the Corn Earworm: Isolation, identification, and synthesis. Science 173: 842-843.
- KING, E. G., G. G. HARTLEY, D. F. MARTIN, J. W. SMITH, T. E. SUMMERS, AND R. D. JACKSON. 1979. Production of the tachnid *Lixophaga diatraea* on its natural

- host, the sugarcane borer, and on an unnatural host, the greater wax moth. Science and Education Administration, Advances in Agricultural Technology. Southern Series. No. 3.
- KING, E. G., AND R. J. COLEMAN. 1989. Potential for biological control of *Heliothis* species. Annu. Rev. Entomol. 34: 53-75.
- KNIPLING, E. F., AND E. A. STADELBACHER. 1983. The rationale for area wide management of *Heliothis* (Lepidoptera: Noctuidae) populations. Bull. Entomol. Soc. America 29: 29-37.
- LEWIS, W. J., AND R. L. BURTON. 1979. Rearing *Microplitis croceipes* in the laboratory with *H. zea* as hosts. J. Econ. Entomol. 63: 656-658.
- PENNACCHIO, F. S., B. VINSON, AND E. TREMBLAY. 1992. In vitro rearing of Cardiochiles nigriceps from egg to second instar. Entomol. Exp. Appl. 64: 209-216.
- POWELL, J. E., AND G. G. HARTLEY. 1987. Rearing *Microplitis croceipes* (Hymenoptera:Braconidae) and other parasitoids of Noctuidae with multicellular host rearing trays. J. Econ. Entomol. 80: 968-971.
- ROTUNDO, G., R. CAVALLORO, AND E. TREMBLEY. 1988. In vitro rearing of Lysiphlebus fabarum (Hym.: Braconidae). Entomophaga 33: 261-267.
- SLANSKY, F. JR. 1986. Nutritional ecology of endoparasitic insects and their hosts: An overview. J. Insect Physiol. 32: 255-261.
- THOMPSON, S. N. 1986. Nutrition and in vivo culture of insect parasitoids. Annu. Rev. Entomol. 31: 197-219.
- Thompson, S. N. 1990. Nutritional considerations in propagation of entomophagous species, pp. 389-404 in New directions in biological control: Alternatives for suppressing agricultural pests and diseases. Alan R. Liss, Inc.
- TILDEN, R. L., AND S. M. FERKOVICH. 1988. Kairomonal stimulation of oviposition into an artificial substrate by the endoparasitoid *Microplitis croceipes* (Hymenoptera: Braconidae). Ann. Entomol. Soc. America 81: 152-156.
- VINSON, S. B. 1975. Biochemical coevolution between parasitoids and their hosts, pp. 14-48 in P. W. Price [ed.]. Evolutionary Strategies of Parasitic Insects and Mites, 225 pp. Plenum, New York.
- WAAGE, J. K., K. P. CARL, N. J. MILLS, AND D. J. GREATHEAD. 1985. Rearing entomophagus insects, pp. 45-66 in P. Singh and R. F. Moore [eds.]. Handbook of Insect Rearing, Vol. 1. Elsevier Press.